



Computer simulation of the isotachophoretic migration and separation of norpseudoephedrine stereoisomers with a free or immobilized neutral chiral selector

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ARTICLE INFO

Article history:

Received 24 February 2020

Revised 24 April 2020

Accepted 27 April 2020

Keywords:

Computer simulation

Electrophoresis

Chiral separation

Electrokinetic chromatography

Isotachophoresis

Cyclodextrin

ABSTRACT

A detailed computer simulation study of the isotachophoretic migration and separation of norpseudoephedrine stereoisomers for cases with the neutral selector added to the leader, immobilized to the capillary wall or support, or partially present in the separation column is presented. The electrophoretic transport of the analytes from the sampling compartment into the separation medium with the selector, the formation of a transient mixed zone, the separation dynamics of the stereoisomers with a free or immobilized selector, the dependence of the leader pH, the ionic mobility of norpseudoephedrine, the complexation constant and selector immobilization on steady-state plateau zone properties, and zone changes occurring during the transition from the chiral environment into a selector free leader are thereby visualized in a hitherto unexplored way. For the case with the selector dissolved in the leading electrolyte, simulation data are compared to those observed in experimental setups with coated fused-silica capillaries that feature minimized electroosmosis and zone detection with conductivity and absorbance detectors.

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1. Introduction

Isotachopheresis (ITP) is performed in a discontinuous buffer system with the sample introduced at the interface between the two electrolytes. Upon current flow, sample components with intermediate effective mobilities compared to those of the electrolyte components of like charge separate according to differences in effective mobilities by forming a pattern of consecutive zones or peaks between the leading zone and the terminating zone with zone properties becoming adjusted according to the electrophoretic regulating principle. The zone pattern attains a migrating steady-state in which all zones have the same velocity (hence the prefix isotacho) and differ in conductivity. Ideally sufficient sample is applied such that zones of constant composition are produced whose length are proportional to the amount of the analyte present. Analytes present in trace amounts become concentrated without forming a plateau-shaped zone and migrate as a sharp peak within a steady-state boundary. ITP analyses can be performed in narrow-bore plastic tubes, separation channels of

rectangular cross section and fused-silica capillaries under conditions with minimized electroosmosis or in presence of electroosmosis. The electrophoretic format can be cationic, anionic or bidirectional [1–15]. In ITP, effective mobilities can be influenced via inclusion of chemical equilibria, including protolysis (proper selection of pH and counter component) [1–15], complex formation between a counter ion and the components to be separated [16,17], complexation of analytes with uncharged or charged additives in the leader [18] or a charged ligand of like charge added to the terminator [19].

Enantioselective separations in capillary electrophoresis are based on the use of chemical equilibria and have been shown to provide high-resolution at low cost for pharmaceutical, pharmacological, agrochemical, environmental, biomedical and forensic analyses [20–25]. Most of the reported techniques are based on capillary zone electrophoresis (CZE) to which one or several chiral selectors, mostly cyclodextrins (CDs), is/are added to the background electrolyte (BGE). Isotachopheretic systems can also be used. However, only relatively few papers can be found in the literature [26–36]. Snopek et al. reported cationic capillary ITP of the enantiomers of various drugs, including pseudoephedrine alkaloids [26] and phenothiazines [27] in presence of neutral CDs. Similarly, the separation of methadone enantiomers, including the isolation of methadone enantiomers by recycling free fluid and con-

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tinuous flow ITP, was reported by our group [28,29]. Dimethindene and pheniramine enantiomers in pharmaceuticals were analyzed by cationic ITP in presence of negatively charged carboxy-ethyl- β -CD [30]. Anionic capillary ITP was employed to separate enantiomers of barbiturates [31], norleucine [32] and tryptophan [33] in presence of neutral CDs. Furthermore, cationic ITP coupled to microcoil NMR detection led to the characterization of enantiomeric separation and interaction between analytes and β -CD [34,35] and inverse cationic chiral ITP configurations in presence of sulfated β -CD provided systems in which either both enantiomers, the enantiomer with weaker complexation or neither of the two enantiomers of a weak base formed ITP zones [36]. In all these cases, chiral separations occur due to the presence of the selector. The technique used is ITP but the separation mechanism is the same as in electrokinetic chromatography [22,23].

Dynamic computer simulation of electrophoretic processes provides insight into particular experimental conditions, including separation mechanisms of analytes. Many dynamic models of various degrees of complexity have been described in the literature [37,38]. The one-dimensional models GENTRANS [39], SIMUL5 [40] and SPRESSO [41] are mostly used and differ in certain aspects but produce identical results when employed with equal input data [42]. The same was recently shown to be true with the electrophoretic transport interface in the COMSOL multiphysics software platform [43]. These approaches do not include chemical equilibria with buffer additives, conversion equilibria of solutes, or solute interactions with column walls or filling material, and can thus not be employed for simulating the separation of enantiomers. Dubrovčáková et al. [44] presented a mathematical model and numerical solution for the addition of a neutral complexation agent to moving boundary systems of strong electrolytes, including the migration of an ITP zone. More recently, GENTRANS [45] and SIMUL5 [46,47] were extended with algorithms that describe 1:1 chemical equilibria between solutes and a buffer additive. These models were found to properly describe the dynamics of chiral separations via use of complexation constants and specific mobilities of formed analyte-selector complexes and were applied to cationic ITP of methadone enantiomers in presence of 2-hydroxypropyl- β -CD [45,48] and sulfated β -CD [36].

ITP has the advantage of regulating the concentration of the analyte in the mM range and thus lends itself to concentrate a solute [1–15]. It can be employed for isolation and identification of single enantiomers in absence of a pure chiral standard and for the preparation of drug enantiomers from racemic mixtures on a micropreparative scale. These aspects are the focus of ongoing research that commenced in our laboratory [28,29,36]. In the present work, two stereoisomers of norpseudoephedrine (NPE), (+)-(1S,2S)-2-amino-1-phenyl-1-propanol ((+)-NPE, also referred to D-NPE or cathine in the literature) and (-)-(1R,2R)-2-amino-1-phenyl-1-propanol ((-)-NPE, also referred to L-NPE in the literature) were used as model compounds. Cathine is found in khat and both NPE stereoisomers are psychostimulant drugs of the phenethylamine and amphetamine chemical classes. They are weak bases. The dynamics of the cationic separation of the NPE stereoisomers was studied in situations with a free or immobilized neutral chiral selector. In analogy to the previous work with methadone [45,48], ITP was performed with sodium and H_3O^+ as leading and terminating components, respectively, and acetic acid as counter component. Heptakis(2,6-di-O-methyl)- β -cyclodextrin (DIMEB) was added as neutral chiral selector. DIMEB was previously shown to resolve the two NPE stereoisomers by ITP [26] and the necessary input constants for simulation were determined by CZE in presence of various amounts of DIMEB [49].

Simulation is shown to provide detailed information about (i) the electrophoretic transport of the analytes from the sampling compartment into the separation medium with a neutral selector

together with the formation of a transient mixed zone, (ii) the ITP separation dynamics of the stereoisomers with a free selector or an immobilized selector, (iii) the dependence of the leader pH, NPE ionic mobility, complexation constant and selector immobilization on steady-state plateau zone properties, and (iv) ITP zone changes occurring during the transition from the chiral environment into a selector free leader solution. Simulation of stereoselective ITP with an immobilized chiral selector and the behavior of ITP zones that migrate from the selector containing region into the leader without chiral selector are presented here for the first time. For the case of ITP in free solution, i.e. ITP with the selector dissolved in the leading electrolyte, simulation data are compared to those observed in experimental setups with coated capillaries that feature minimized electroosmotic flow (EOF) and zone detection with conductivity and absorbance detectors placed along the column.

2. Materials and Methods

2.1. Computer simulations

Simulations were performed with GENTRANS which comprises algorithms to account for the interaction with an electrolyte additive [45,48]. It is assumed that these equilibria are instantaneous which means that the kinetics of complex formation do not play a considerable role during migration. In addition to the electrophoretic mobility and the pK_a values of each component in the system, complexation constants and the mobilities of the analyte-additive complexes are used as input. The model does not take into account the dependence of mobilities, pK_a values and complexation constants on ionic strength, viscosity and temperature. The program was executed on Windows XP or 32bit Windows 7 based PC's featuring Intel Core2 Duo 2.93 GHz and Intel Core i5 2.8 GHz processors, respectively. If not stated otherwise, a 10 cm column divided into 20000 segments (5 μm mesh) with the sample being placed between 5 and 6 % of column length (boundary width: 0.001 %) was assumed, a constant current density of 1000 A/m² was applied and EOF was omitted. Simulations for 2.5 min electrophoresis time lasted 16–20 h. The component's input data for simulation are summarized in Table 1. Only NPE stereoisomers were assumed to interact with DIMEB. The complexation constants and mobilities for NPE stereoisomers listed in Table 1 are those determined previously by chiral CZE in a pH 4.10 acetic acid buffer (47 mM acetic acid adjusted with NaOH to pH 4.10) [49]. For making plots, simulation data were imported into SigmaPlot Scientific Graphing Software version 12.5 (Systat Software, San Jose, CA, USA).

2.2. Chemicals and samples

All chemicals used were of analytical or research grade. (+)-NPE and (-)-NPE as hydrochloride salts were from Fluka (Buchs, Switzerland) whereas the free bases were purchased from Sigma Aldrich (Buchs, Switzerland). Acetic acid was supplied from Merck (Darmstadt, Germany). DIMEB was from Cyclolab (Budapest, Hungary). The pH of the leading electrolyte (catholyte) was adjusted with 0.1 M acetic acid. Samples were prepared in water and did not contain a chiral selector.

2.3. Electrophoretic instrumentation for chiral ITP and running conditions

The ITP separation process of NPE stereoisomers was monitored with a laboratory made system comprising a 50 μm I.D. linear polyacrylamide (LPA) coated fused-silica capillary of 70 cm length (Polymicro Technologies, Phoenix, AZ, USA), a purpose built sequential injection analysis manifold for fluid handling and sample

Table 1
Physico-chemical input parameters used for simulation

Compound	pK _a	Mobility (10 ⁻⁸ m ² /Vs)	Complex constant (L/mol)	Complex mobility (10 ⁻⁸ m ² /Vs)	Ref.
Sodium	-	5.19	-	-	c
Acetic acid	4.76	4.24	-	-	c
Chloride	-	7.12	-	-	c
(-)-NPE	8.9	2.65	44.7	0.442	[49] ^b
(+)-NPE	8.9	2.65	66.5	0.554	[49] ^b
DIMEB	^a	1.00	-	-	

^a With GENTRANS, DIMEB as neutral cyclodextrin is entered as a weak acid with a pK_a of 14.^b Complex constant and complex mobility values are those for complexation of the protonated bases with DIMEB in a 47 mM acetate buffer at pH 4.10 [49]^c From database of mobilities and pK_a values of SIMUL5 [40].

injection and an array of 8 contactless conductivity detectors as is described in detail elsewhere [50]. The LPA coated capillary exhibits a low EOF (mobility of 2.34×10^{-9} m²/Vs) and the manifold allows the placement of the first detector at about 2.7 cm from the sample inlet. The centers of the 8 detectors were positioned 2.7, 8.2, 13.2, 18.4, 23.5, 28.5, 33.5 and 38.5 cm away from the capillary's injection end (distance between detectors was about 5 cm). Other ITP experiments were made with an autosampler PrinCE-C 560 2-Lift (Prince Technologies, Emmen, The Netherlands) at ambient temperature (about 25°C) using an LPA coated fused-silica capillary of 50 μ m I.D. and 83 cm total length (Polymicro Technologies) and two detectors, a UVIS 206 PHD detector (Linear Instruments, Reno, NV, USA) operated at 200 nm and placed at 50.5 cm (60.8 % of column length) and a TraceDec contactless conductivity detector (Innovative Sensor Technologies, Strasshof, Austria) at 61.4 cm (74.0 % of column length). This setup was previously used for chiral ITP [48] and CZE [49]. The PrinCE autosampler was also used with a permanently coated Guarant capillary (Alcor BioSeparations, Palo Alto, CA, USA; electroosmotic mobility of 1.23×10^{-9} m²/Vs) of 50 μ m I.D. and 70 cm total length and two conductivity detectors, a laboratory made high voltage contactless conductivity detector (provided by Dr. Peter Hauser, University of Basel, Basel, Switzerland) at about 33.4 cm and a TraceDec (Innovative Sensor Technologies) detector at about 46.3 cm. This setup was previously used for chiral ITP of methadone [48]. Both types of the coated capillaries used were conditioned using water and leading electrolyte. Between runs, the capillaries were rinsed with water, 70% 2-propanol and water. Before each run the capillary was rinsed with leading electrolyte, the sample was applied at the anodic end via application of pressure, and the anodic end of the capillary was dipped into the anolyte. A constant current of 2.0 or 3.0 μ A, or constant voltage of 20 kV (current < 6 μ A), was applied. Detector data were collected and stored using a 16-channel (array detector setup) or a 4-channel (dual detector setups) e-corder (eDAQ, Denistone East, NSW, Australia).

3. Results and discussions

3.1. Isotachopheresis of NPE stereoisomers

NPE is a weak base that migrates isotachophoretically with sodium as leading constituent, H₃O⁺ or β -ala as terminating ion and acetic acid as counter component [26,50,51]. Enantioseparation is achieved in presence of DIMEB. Simulation data presented in Fig. 1 were obtained with a leading electrolyte (catholyte) composed of 10 mM NaOH, 24.6 mM acetic acid (pH 4.60) and 10 mM DIMEB. 10 mM acetic acid (pH 3.39) with 10 mM DIMEB served as terminating electrolyte (anolyte). The sample was composed of 2.85 mM of each NPE base and did not contain other components. With the interaction between NPE and DIMEB, simulation predicts a separation of the NPE stereoisomers between sodium and the Kohlrausch adjusted acetic acid solution (Fig. 1A). (-)-NPE with the lower complexation constant is forming an isotachophoretic zone

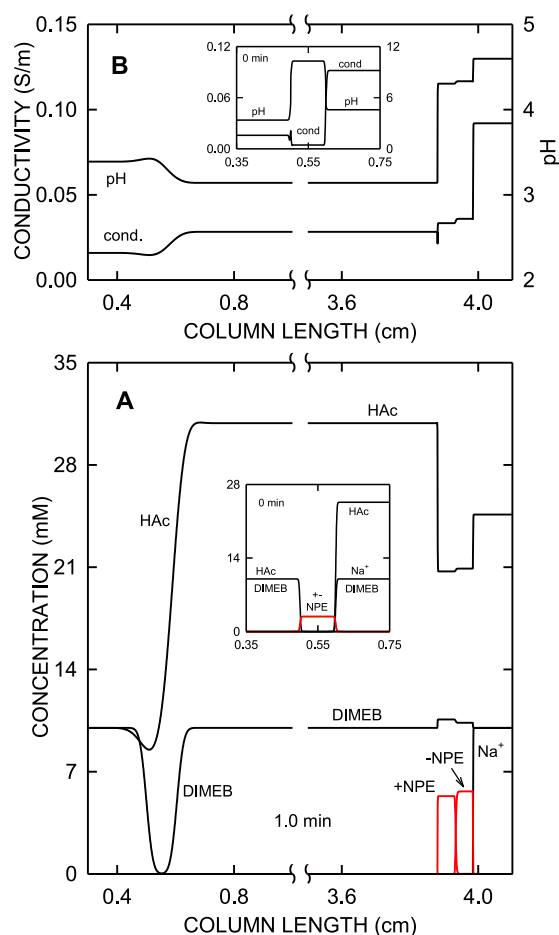


Figure 1. Computer predicted separation of NPE stereoisomers utilizing a pH 4.60 leading electrolyte (catholyte) composed of 10.0 mM NaOH, 24.6 mM acetic acid and 10 mM DIMEB. A mixture of 10.0 mM acetic acid and 10 mM DIMEB served as terminating electrolyte (anolyte). The sample comprised (+)-NPE and (-)-NPE as bases (2.85 mM each) without any other compounds. The simulation was performed at a constant current density of 1000 A/m² and without any EOF. (A) Computer predicted NPE (red line graphs), acetic acid, sodium and DIMEB concentration profiles after 1 min and initial distributions (insert). (B) pH and conductivity profiles after 1 min and initial distributions (insert). -NPE and +NPE refer to (-)-NPE and (+)-NPE, respectively.

between sodium, the leading ion, and (+)-NPE because its net mobility is larger compared to that of (+)-NPE and smaller than that of sodium. (+)-NPE produces an isotachophoretic zone which migrates between that of (-)-NPE and the adjusted acetic acid zone with H₃O⁺ as terminating ion. This is comparable to the behavior of the methadone enantiomers in presence of hydroxypropyl- β -CD as reported previously [45,48]. The concentration of acetic acid in the adjusted terminating zone (30.87 mM) is higher compared to that in the leader (24.6 mM) and the 10 mM applied as anolyte

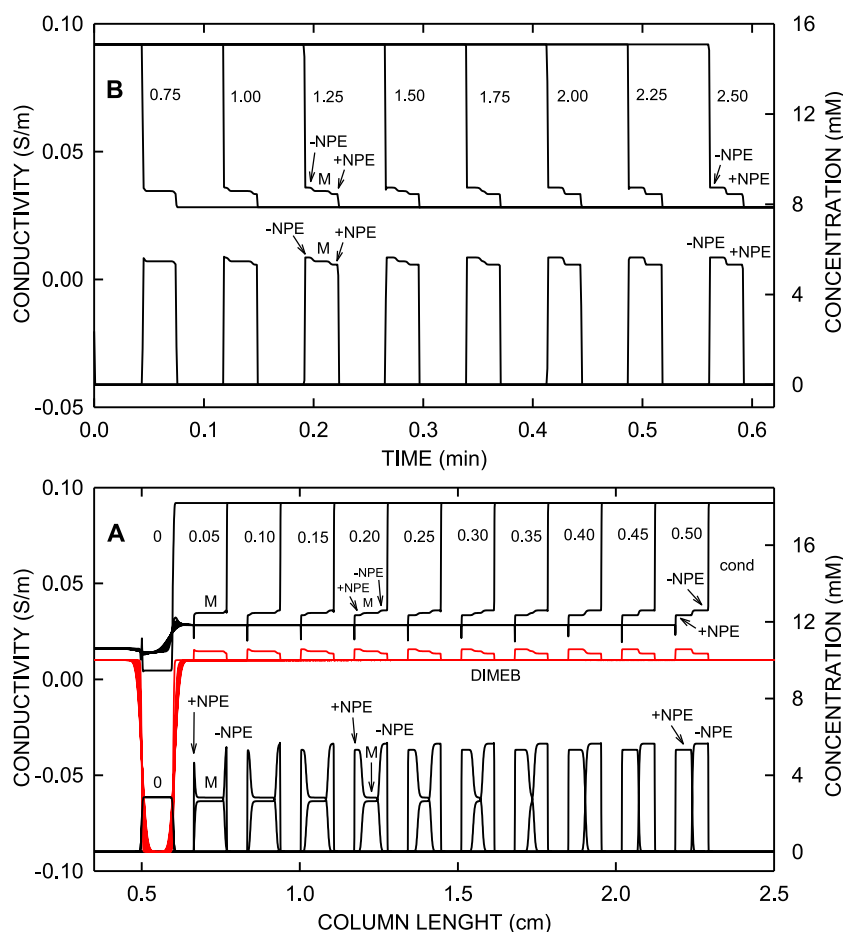


Figure 2. Computer predicted (A) analyte profiles (lower graphs), DIMEB distributions (red line graphs) and conductivity profiles (upper graphs) at 0, 0.05, 0.10, 0.15, 0.20, 0.25, 0.30, 0.35, 0.40, 0.45 and 0.50 min of power application, and (B) detector profiles for absorbance (lower graphs) and conductivity (upper graphs) for detectors placed at 0.75, 1.00, 1.25, 1.75, 2.00, 2.25 and 2.50 cm of column length. Simulation conditions as for Fig. 1. -NPE, +NPE and M refer to (-)-NPE, (+)-NPE and mixed zone, respectively.

(Fig. 1A). Furthermore, zone conductivities and pH values of the NPE zones are predicted to be between those of the adjusted terminating zone (28.3 mS/m; 3.14) and the leader (92.0 mS/m, 4.60) (Fig. 1B). The NPE zones are characterized with sharp front and rear boundaries and the rear boundary features a conductivity dip which is comparable to previously described cationic ITP configurations with H_3O^+ as terminating ion [45,48].

The formed (-)-NPE and (+)-NPE sample zones not only differ in the plateau concentration (5.64 vs. 5.33 mM, respectively), pH (4.33 vs. 4.31), conductivity (36.0 vs. 33.4 mS/m), acetic acid concentration (20.9 vs. 20.7 mM) and ionic strength (5.69 vs. 5.38 mM), but also in the concentration of DIMEB (10.35 vs. 10.57 mM). DIMEB is neutral and does not migrate under the influence of the electric field. The DIMEB increase inside the ITP zones compared to its value in the leading and terminating electrolytes is due to the migration of the charged complexes and was previously reported for the interaction of a neutral cyclodextrin with strong electrolytes [44] and the weak base methadone [48]. The increase of a neutral CD within the ITP zones of various pharmaceutical compounds could be experimentally visualized with on-line microcoil NMR detection [52]. For migrating analytes in CZE that interact with the selector, CD concentration deviations from its value in the buffer were also predicted with dynamic simulation [45] and a generalized model of the linear theory of electromigration [49,53]. An increase of CD inside the migrating analyte zone could be experimentally validated using a neutral CD with absorbance at 245 nm together with a non-absorbing analyte [53]. In absence of complexation, there is no DIMEB change and no separation, and the

adjusted total NPE concentration in the investigated ITP system is 6.84 mM. For the data of Fig. 1 obtained with a 10 cm column and a constant $1000 A/m^2$, the predicted voltage increased from the initial 1544 V to 1806, 2218 and 3456 V within 0.5, 1.0 and 2.5 min of current flow, respectively.

3.2. Separation process

Separation in ITP proceeds via migrating transient mixed zones which are formed according to the regulating principle [9–15]. For the example of Fig. 1, this is illustrated with the simulation data presented in Fig. 2. The properties of the mixed zone, including its conductivity, are distinctly different to the properties of the pure zones of (-)-NPE and (+)-NPE which are gradually formed in front of and behind the mixed zone, respectively (Fig. 2A). The mixed zone becomes smaller with time and vanishes when the separation of the two stereoisomers is completed (lower graphs in Fig. 2A). For the presented case, separation is predicted to become complete shortly after 0.4 min of current application (Fig. 2A). At 0.4 min the boundary between the stereoisomers has not yet reached a steady-state shape. It is distinctly broader compared to the boundary predicted after 0.45 min and 0.50 min. Once steady-state is reached, the separated stereoisomers continue to migrate as a steady-state migrating zone pattern.

The conductivity (34.6 mS/m) and the DIMEB concentration (10.47 mM) of the mixed zone are predicted to be between the conductivities and DIMEB concentrations of the pure zones (red and upper graphs in Fig. 2A). The separation can be followed with

multiple detectors placed along the separation column. The simulation data presented in Fig. 2B were generated for UV absorbance (lower graphs representing the sum of the NPE concentrations) and conductivity (upper graphs) with detectors placed at 0.75, 1.00, 1.25, 1.75, 2.00, 2.25 and 2.50 cm of column length (from left to right, respectively). Data storage occurred at 20 Hz. It is important to note that this frequency was too low to monitor the sharp conductivity dip at the interface between (+)-NPE and the adjusted terminator zone (see conductivity profiles in Fig. 1B and 2A). It enabled, however, the prediction of the transient mixed zone and the formation of the two NPE ITP zones as would be observed by absorbance and conductivity detection along the column.

For validation, the separation of (-)-NPE and (+)-NPE was monitored with a purpose made instrument featuring 8 contactless conductivity detectors along a 70 cm LPA coated capillary which exhibits a low EOF towards the cathode. This approach was previously employed to study the formation of the leader/terminator boundary in absence of the sample as well as the formation of an ITP zone of NPE in absence of the chiral selector [50]. In analogy to these efforts, the separation of the two stereoisomers was simulated with a 17.5 cm separation space divided into 35000 segments of equal length ($\Delta x = 5.0 \mu\text{m}$) and a constant current density of 370 A/m^2 . Sample was applied at the anodic capillary end as a plug with a length of 2.5 % of column length (initially placed between 5.0 and 7.5 % of column length) and a boundary width of 0.02 % (for distribution, see lower insert in Fig. 3A). For 10 min of electrophoresis, this required a simulation time of about 30 h. The compositions of the leading and terminating electrolytes as well as that of the sample were the same as for the simulation example presented in Fig. 1 and the input data used were those of Table 1. Furthermore, in order to mimic the temporal behavior of the EOF present in the experiment, the ionic strength dependent EOF model employed previously for this ITP system without chiral selector (for details refer to [50]) was applied. The predicted detector profiles for the 8 detectors (4 Hz data that do not show the conductivity dip) are depicted in Fig. 3A. They illustrate the expected separation process as discussed with the data of Fig. 2A and 2B. The voltage is predicted to increase from 1056 to 2543 V within the 10 min time interval (solid line graph in upper insert of Fig. 3A). The EOF increases from 25.5 to $64.7 \mu\text{m/s}$ (dashed line graph in upper insert of Fig. 3A) and is significantly smaller compared to the electrophoretic transport rate of $211.0 \mu\text{m/s}$ reported in Ref. [50]. The predicted net transport thus increases from 236.5 to $275.7 \mu\text{m/s}$.

Experimental data obtained under a constant current of $3 \mu\text{A}$ are presented in Fig. 3B. The existence of the mixed zone and the gradual decrease of its length could thereby be experimentally verified. Experimentally monitored detector profiles of detectors 2 to 7 correspond qualitatively well with those predicted by simulation. For the chosen conditions under constant current density, the voltage is predicted to increase in a similar fashion as was predicted by simulation (compare voltage graphs in upper insert of Fig. 3A with insert of Fig. 3B).

For the first detector (D1 of Fig. 3) and the transition marked with asterisk detected with the second detector (D2), the predicted detector profiles are somewhat different compared to those monitored experimentally. It was previously noted that sample injection in the employed SIA setup did not provide sharp initial boundaries [50]. Broader initial boundaries between sample and its surrounding electrolytes (L and T on cathodic and anodic side, respectively) were found to have an impact on the predicted detector signal of the first detector and the buffer transition of the second detector marked with an asterisk in Fig. 3. This is illustrated with the data of Fig. 3C which were obtained with dispersed initial sample boundaries (lower inset in Fig. 3C) which were produced from the initial distribution of Fig. 3A via application of Taylor-Aris disper-

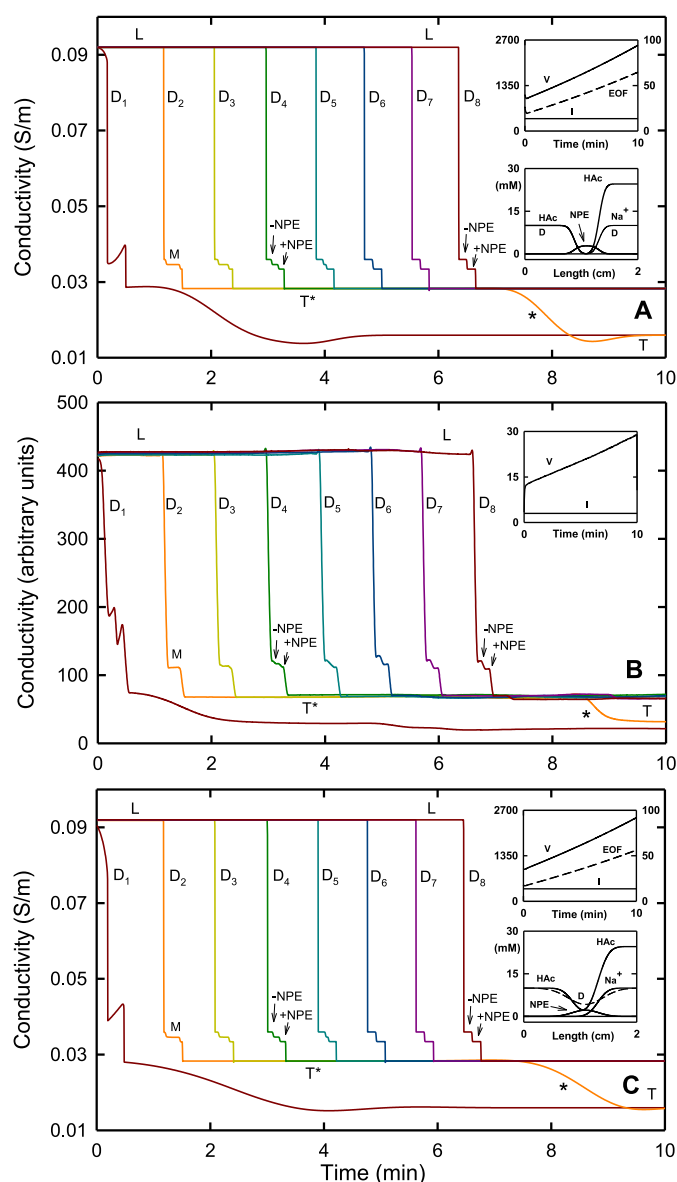


Figure 3. (A,C) Simulated and (B) experimental conductivity electropherograms of detectors D1 to D8 for detection of the NPE stereoisomer separation under constant current conditions in an LPA coated fused-silica capillary. The leading electrolyte (L, catholyte) was composed of 10 mM NaOH, 24.6 mM acetic acid and 10 mM DIMEB. The terminating electrolyte (T, anolyte) was 10 mM acetic acid. Simulations were performed with an ionic strength dependent EOF model using (A) sample boundaries with a width of 0.02 % and (C) dispersed initial sample boundaries. For details refer to text. The inserts in panels A and C comprise the initial distributions (lower graphs, with D for DIMEB) and the computer predicted temporal behavior of voltage V (V, left y-axis scale), current density I (A/m^2 , left y-axis) and EOF ($\mu\text{m/s}$, right y-axis scale) depicted as upper graphs. The insert in panel B represents recorded current (μA) and voltage (kV). The asterisk marks the transition between adjusted terminator (T^*) and the terminating electrolyte (T) and represents the EOF marker.

sion in a $50 \mu\text{m}$ ID capillary at a flow rate of 1.5 mm/s for 0.5 min (for impact of flow on dispersion refer to [54]). Better agreement between experimental and simulation data for detector D1 and for the buffer transition of detector D2 was thereby predicted.

3.3. Formation and composition of the mixed zone

For the example presented in Fig. 2, the mixed zone is characterized with a higher concentration of (+)-NPE (2.83 mM), the stereoisomer which has a stronger interaction with DIMEB, compared to (-)-NPE (2.64 mM). This is comparable to the separation

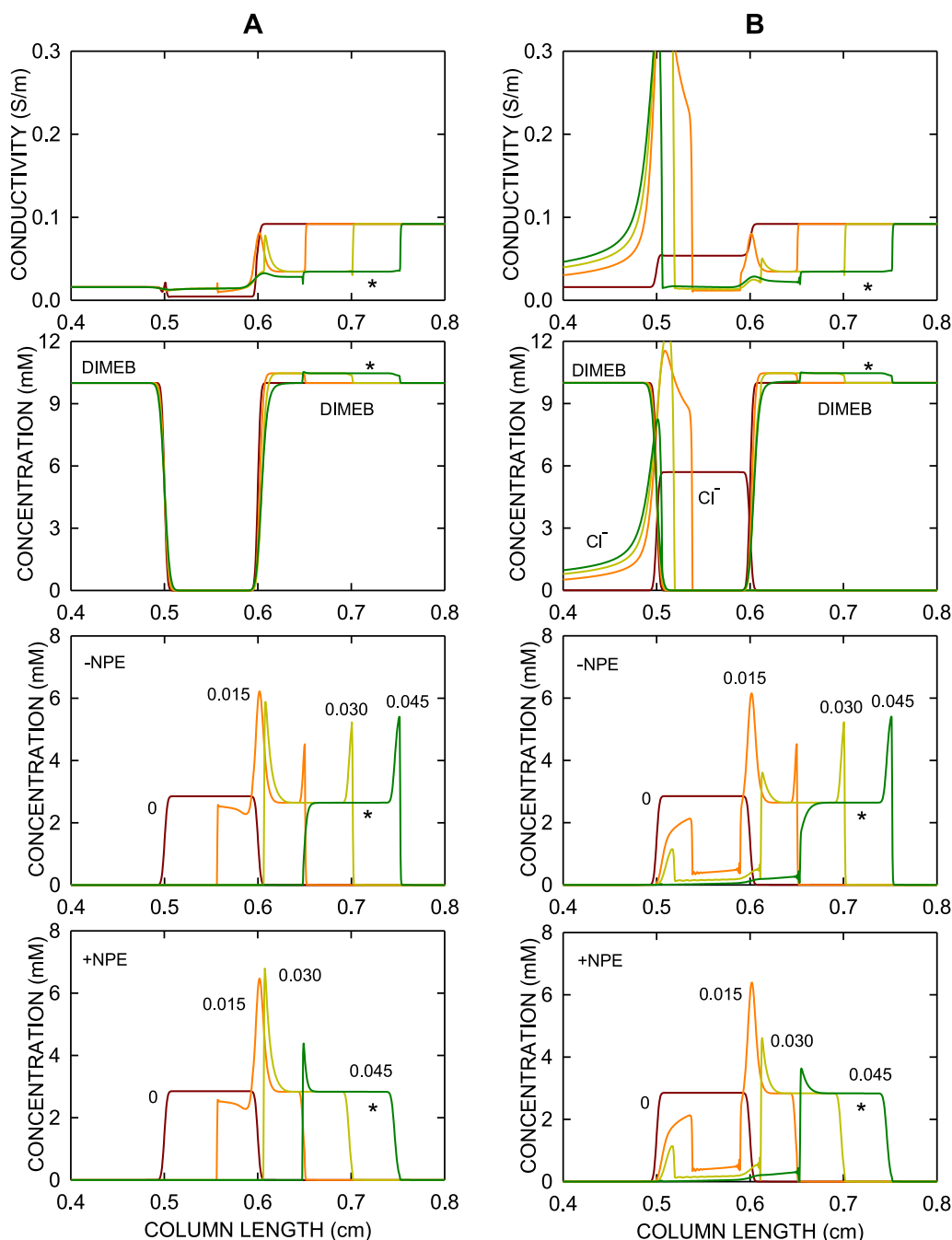


Figure 4. Computer predicted concentrations of (+)-NPE (bottom panels), (-)-NPE (second panels from bottom), DIMEB and chloride (second panels from top) and conductivity distributions (top panels) for NPE analytes sampled (A) as free bases and (B) as hydrochlorides. Profiles shown are for 0 min (dark red graphs), 0.015 min (orange graphs), 0.030 min (yellow graphs) and 0.045 min (green graphs) of current flow. The asterisks mark the property levels of the mixed zone. Other conditions as for Fig. 1 and 2.

of methadone enantiomers described previously [48] and appears to be typical for separations in which complexation with a neutral CD is involved. It is important to note that with an equimolar mixture of two weak bases and separation of these bases in absence of complexation, an equimolar mixed zone would be formed.

Simulation also revealed that the composition of the formed mixed zone is independent of the sample composition. This is illustrated with the concentration and conductivity data presented in Fig. 4. Profiles of the initial distributions and those after 0.015, 0.030 and 0.045 min (0.9, 1.8 and 2.7 s, respectively) of current flow at a current density of 1000 A/m² are depicted. These data illustrate the transients occurring after application of power and the formation of the mixed zone (property levels marked with

asterisks in Fig. 4) outside the sampling section. Data presented are those for the two NPE stereoisomers sampled as weak bases (Fig. 4A, case of Fig. 1 and 2 with 2.85 mM of each) and as hydrochlorides (Fig. 4B, 2.85 mM of each stereoisomer together with 5.7 mM chloride). Note that the samples did not contain buffer components and DIMEB, and were applied between 0.5 and 0.6 cm.

Upon current flow, the transition from the sampling to the separation column (at 0.6 cm) is characterized with transient NPE peaks that occur at the cathodic interface with the leader where NPE becomes complexed with DIMEB and where NPE reaches a region of higher conductivity (lower electric field strength). It is also the location where the migrating steady-state front boundary is formed due to the cathodic migration of the leading ion (Na⁺).

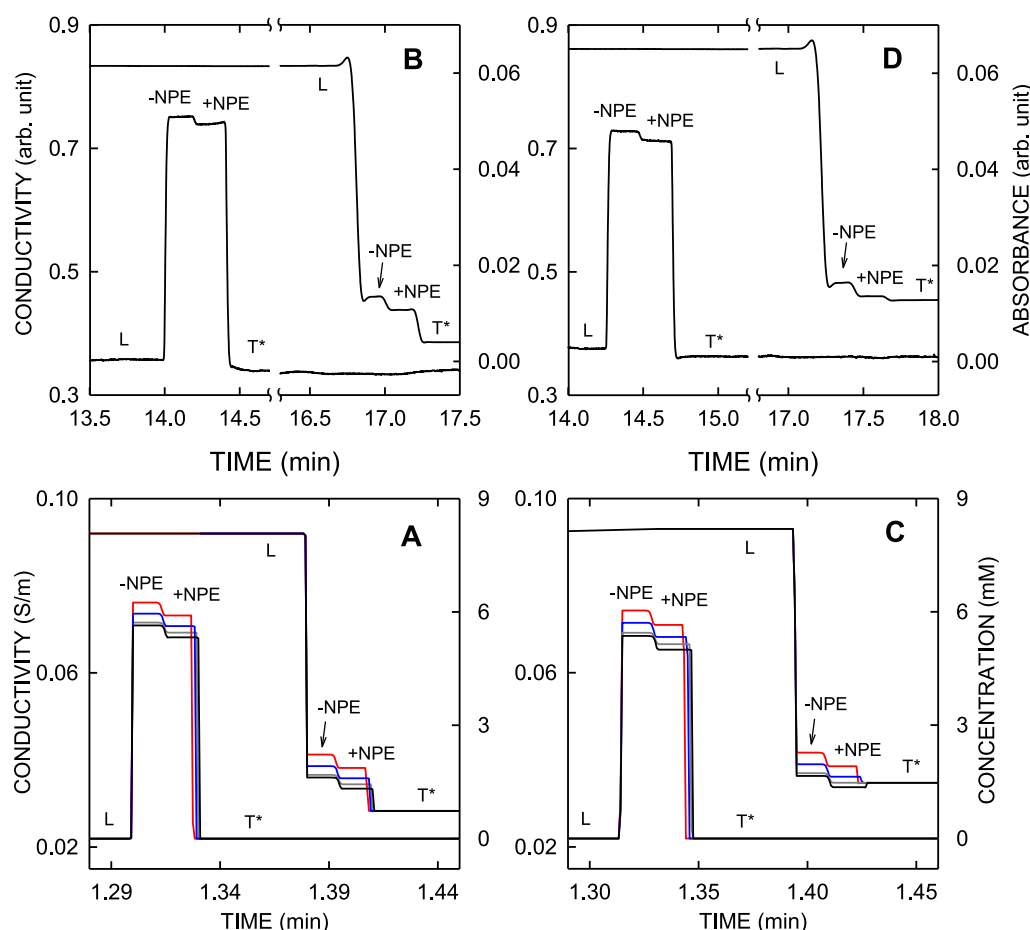


Figure 5. (A,C) Computer predicted detector profiles for a detector placed at 5 cm of column length measuring absorbance (sum of NPE concentrations) and conductivity (shifted by 0.08 min for presentation purposes) and (B,D) experimental data obtained with the setup featuring an LPA capillary with absorbance (200 nm) and conductivity detection for ITP systems with leading electrolyte pH values of (A,B) 4.60 and (C,D) 4.30. Simulation and experimental data were obtained with NPE stereoisomers (2.85 mM each) sampled as hydrochlorides. Other simulation conditions as described in the text. In the experiments, sample application occurred at 80 mbar for 0.3 min and the run was performed at a constant 2 μ A (voltage change between about 3 and 21 kV). -NPE, +NPE, L and T* refer to (-)-NPE, (+)-NPE, leading electrolyte and adjusted terminator, respectively.

In absence of chloride in the sample, NPE is predicted to completely migrate from the sampling compartment into the separation column within 0.030 min (Fig. 4A). Due to chloride, which migrates in the opposite direction and strongly influences local conductivity, NPE requires more than 0.045 min to become completely transported into the separation column (Fig. 4B). After about 0.15 min of current flow, profiles of (+)-NPE, (-)-NPE and the mixed zone of the two investigated cases become indistinguishable (data not shown). These data illustrate how simulation can be used to provide insight into the processes occurring due to electrophoretic transport of a given electrolyte distribution.

3.4. Impact of input data and comparison to experimental results

Simulated detector profiles of the isotachophoretic zones of (-)-NPE and (+)-NPE predicted for detection with a UV absorbance detector and a conductivity detector are presented in Fig. 5A. The data correspond to a detector location at 5 cm of column length (center of the column; 20 Hz signal) and the conditions of Fig. 1 with a leader pH of 4.6. The absorbance data represent the sum of the NPE concentrations. For presentation purposes, the conductivity data are depicted with a time shift of 0.08 min. Black line graphs are obtained with the input data of Table 1. These data are in qualitative agreement with those monitored experimentally using the setup with an LPA coated fused-silica capillary interfaced

with a UV absorbance and a conductivity detector that were placed about 11 cm apart (Fig. 5B, cf. Section 2.3). In an effort to find optimized input data for ITP, the mobility of the free NPE molecule was varied. An increase in the free mobility value resulted in increased NPE concentrations in the formed ITP zones and thus somewhat shorter zones. This is illustrated with the simulated absorbance data for mobilities of 2.65×10^{-8} m²/Vs (black graph), 2.85×10^{-8} m²/Vs (blue graph) and 3.05×10^{-8} m²/Vs (red graph) presented in Fig. 5A.

Best agreement between computer predicted and measured conductivity distributions was obtained using an NPE free mobility of 2.85×10^{-8} m²/Vs (blue graph in Fig. 5A) instead of the 2.65×10^{-8} m²/Vs (black graph in Fig. 5A). A free mobility of 3.05×10^{-8} m²/Vs resulted in data with a too high conductivity within the NPE zones (red graph in Fig. 5A). Variation of the input data for complexation of NPE with DIMEB has only a minor impact on the distributions of conductivity and other properties and is in analogy to the case of methadone discussed previously [48]. For NPE, this is illustrated with the gray line graph of Fig. 5A which was obtained with a different set of input data. The complexation constants for (-)-NPE and (+)-NPE were 54.4 L/mol and 76.3 L/mol, respectively, and the complex mobilities 0.853×10^{-8} m²/Vs and 0.887×10^{-8} m²/Vs, respectively. The free mobility of NPE used for this simulation was that listed in Table 1. These data illustrate that the mobility value determined by CZE at a pH 4.10 and an ionic

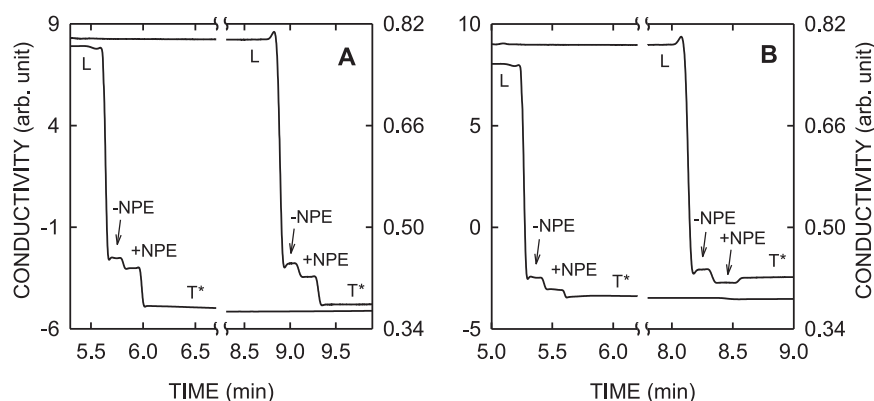


Figure 6. Experimental data of separated NPE stereoisomers having leading electrolyte pH values of (A) 4.60 and (B) 4.30 obtained in the setup with a Guarant capillary and two conductivity detectors (y-axis scales of first and second detector are on the left and on the right, respectively). Sample application occurred at 80 mbar for 0.3 min and experiments were performed at a constant 20 kV. Key as for Fig. 5.

strength of 8.58 mM [49] are close but somewhat too low for the conditions encountered in ITP. Temperature or ionic strength differences might contribute to this fact.

In order to gain further insight into the validity of a higher free mobility for NPE, the ITP system was changed by lowering the pH of the leader to 4.30. Simulation data obtained with a leader composed of 10 mM NaOH, 40 mM acetic acid and 10 mM DIMEB are presented in Fig. 5C and corresponding experimental data are depicted in Fig. 5D. As is the case with a leader pH of 4.60 (Fig. 5A and 5B), a mobility of $2.85 \times 10^{-8} \text{ m}^2/\text{Vs}$ (blue graph in Fig. 5C) provides a good match with the experimental conductivity data. With a mobility of $2.65 \times 10^{-8} \text{ m}^2/\text{Vs}$ (black graph in Fig. 5C), the predicted conductivities of the two NPE zones are lower compared to those monitored experimentally. The calculated conductivity of the (+)-NPE zone is even lower compared to that of the adjusted terminating zone (zone T*). With a mobility of $3.05 \times 10^{-8} \text{ m}^2/\text{Vs}$ (red graph in Fig. 5C), the predicted conductivities of the two NPE zones are higher compared to those measured experimentally. These data illustrate that a free NPE mobility of $2.85 \times 10^{-8} \text{ m}^2/\text{Vs}$ provides simulation data that compare well with experimental isotachopherograms obtained at different pH values of the leader.

In addition to the experimental setup with an LPA coated capillary together with absorbance and conductivity zone detection (Fig. 5B and 5D), a setup with a permanently coated Guarant capillary and two conductivity detectors placed about 13 cm apart (cf. Section 2.3) was used to follow and characterize the NPE ITP zones in presence of 10 mM DIMEB and application of constant voltage instead of constant current. The second conductivity detector is the same as that employed together with absorbance detection. Data obtained with leader pH values of 4.6 and 4.3 are presented in Fig. 6A and 6B, respectively. In both cases, the conductivities of the NPE zones detected with the second detector were lower compared to those monitored with the first detector. The same was found to be true for the case of methadone enantiomer detection reported previously [48]. The reason for this change is unknown and was not further investigated. It is interesting to mention that such a change was not observed with the conductivity array detector under both constant current (Fig. 3B) and constant voltage (data not shown) conditions.

3.5. Immobilization of the chiral selector

Neutral CDs can also be immobilized, i.e. bound to the inner wall of a capillary, to particles used as capillary packing material, to a monolith present in the capillary, or to nanoparticles which are dispersed in the running buffer. These approaches are

referred to as open-tubular, packed, monolithic and pseudostationary phase capillary electrochromatography (CEC), respectively, and have been applied to separations in uniform background electrolytes [56,57]. In analogy to the CEC simulations in a uniform buffer presented previously for the separation of enantiomers of a weak base [55], computer simulation was employed to study the isotachopheretic migration of (+)-NPE and (-)-NPE in presence of immobilized DIMEB. The impact of the 1:1 interaction between NPE and DIMEB was simulated by setting the complex mobilities and the mobility of DIMEB (which is used for the calculation of the diffusion coefficient) to zero. All other input parameters listed in Table 1, including the complexation constants, were assumed to be identical to those in free solution. This provides data that mimic migration and separation with the chiral selector being immobilized to the capillary wall or support material and without unspecific interactions between analytes and the chiral stationary phase. Data predicted for the system of Fig. 1 with immobilized DIMEB are presented in Fig. 7.

The simulation data given in Fig. 7 reveal that the formation of ITP zones of the NPE stereoisomers are predicted also for the case of DIMEB immobilization. This is the first account for ITP of stereoisomers in presence of an immobilized selector. The data presented in Fig. 7A indicate that the separation of (+)-NPE and (-)-NPE proceeds in the same manner as is the case for free solution (compare with Fig. 2A). Separation, however, is predicted to be faster when DIMEB is immobilized (about 0.30 min with immobilized DIMEB vs. about 0.45 min in the case of Fig. 2A). The electrophoretic displacement rates are identical as this transport rate is determined by the components of the leading electrolyte, i.e. compounds that were assumed not to interact with DIMEB, and the current density [50]. Acetic acid was reported to exhibit a very weak interaction with DIMEB [58] which does not have an appreciable impact on the displacement rate and the formation of the migrating NPE zone (simulation data not shown).

All properties of the formed ITP zones are somewhat lower compared to those predicted for the case of free solution. The plateau concentrations of (-)-NPE and (+)-NPE are 5.40 and 4.91 mM, respectively (5.64 and 5.33 mM for free solution), the pH values are 4.31 and 4.26, respectively (4.33 and 4.31), the conductivities are 34.02 and 30.32 mS/m, respectively (36.0 and 33.4 mS/m) and the acetic acid concentrations are 20.75 and 20.49 mM, respectively (20.9 and 20.7 mM). The concentration of DIMEB remains 10 mM (no change because DIMEB is immobilized, Fig. 7A and 7B) which is in contrast to the case of free solution (10.35 and 10.57 mM). Furthermore, analyte transition between sample compartment and separation column is essentially identical to that described in Fig. 4A (data not shown). Experimental validation of the

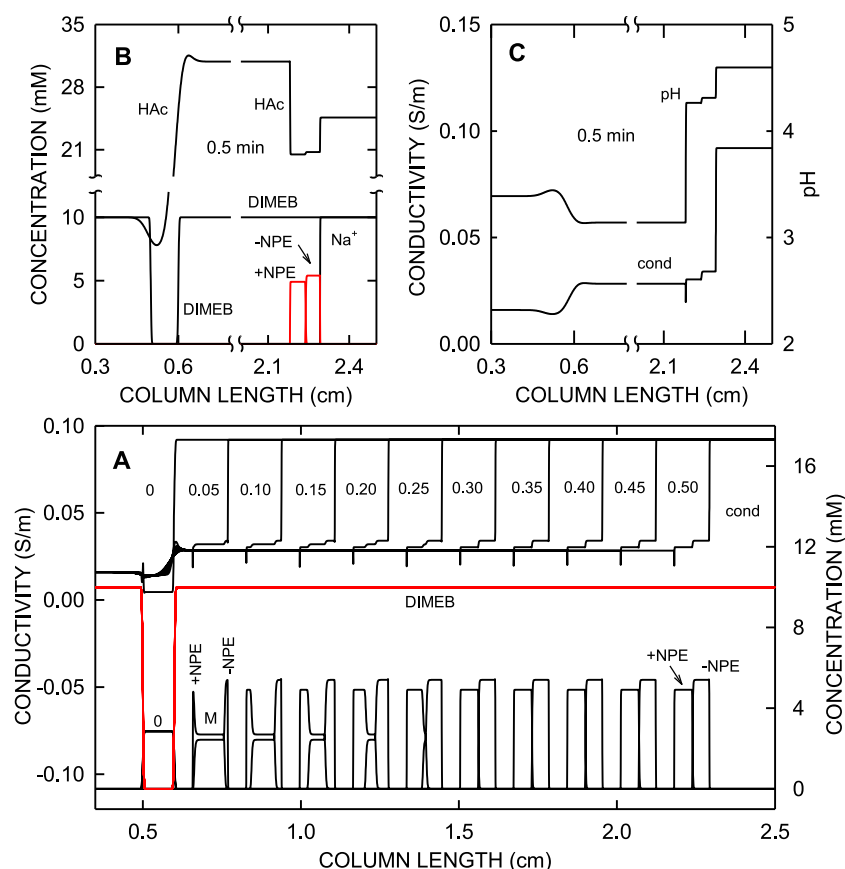


Figure 7. Computer predicted NPE stereoisomer separation in presence of 10 mM immobilized DIMEB along the separation column and otherwise identical conditions as for Fig. 1. For details refer to text. (A) Analyte profiles (lower graphs), DIMEB distribution (red line graph) and conductivity profiles (upper graphs) at 0, 0.05, 0.10, 0.15, 0.20, 0.25, 0.30, 0.35, 0.40, 0.45 and 0.50 min of application of a constant 1000 A/m². (B) Concentration distributions of all components (NPE stereoisomers as red line graphs) after 0.5 min. (C) Conductivity and pH profiles after 0.5 min. Key as for Figs. 1 and 2.

predicted zone structure, e.g. via open-tubular CEC, remains to be undertaken.

3.6. Partial filling of column with chiral selector

Simulation was also used to study the analyte behavior at the rear end of a DIMEB zone, i.e. a situation with partial filling of the column with DIMEB. This includes a region where ITP zones leave the environment with complexation, is relevant for off-column analyte detection, e.g. with mass spectrometry (MS), and is described here for the first time via the dynamics of migrating analyte zones. Simulation data for a column with DIMEB up to 5 cm of column length, a sample composed of NPE stereoisomers applied as hydrochlorides, and otherwise identical column conditions as for Fig. 1 are presented in Fig. 8. Within the DIMEB segment, (-)-NPE and (+)-NPE were predicted to become separated within about 0.45 min and to migrate isotachophoretically as consecutive zones up to the end of the segment with DIMEB (Fig. 8). The front boundary of the (-)-NPE ITP zone reached the vanishing DIMEB gradient shortly after 1.25 min. Upon leaving complexation, both stereoisomers become concentrated to 6.84 mM and jointly form a migrating ITP zone of uniform conductivity (46.97 mS/m, see 1.50 min time point of Fig. 8). Acetate concentration and pH within this zone are also uniform (21.78 mM and 4.42, respectively; profiles not shown). This ITP zone exhibits a non-uniform distribution of the two analytes. (-)-NPE is still migrating ahead of (+)-NPE and the two stereoisomers will become mixed with time (Fig. 9A). It is important to realize that the migration rate of the newly formed ITP zone remains equal to that of the pattern within the DIMEB

zone. It is noteworthy to mention that DIMEB is present in solution. It has a higher value than 10 mM in the ITP zones (see 0.25 to 1.25 min profiles depicted as red line graphs of Fig. 8) and the mixed zone (see 0.25 min profile of Fig. 8). As a result of diffusion the DIMEB gradients at 5 cm of column length and at the edges of the sample compartment (0.5 and 0.6 % of column length) broaden with time.

The processes occurring during the migration of the ITP zones across the DIMEB boundary at 5 cm of column length and thereafter is depicted in Fig. 9. It represents a situation in which DIMEB is immobilized with a boundary width of 0.01 %. DIMEB is therefore invariant and present as non-diffusing gradient at its zone edge. Within the immobilized DIMEB segment, the NPE stereoisomers were predicted to become separated within about 0.30 min (Fig. 7A) and to migrate isotachophoretically as consecutive zones up to the end of the segment with immobilized DIMEB (Fig. 9A). The front boundary of the (-)-NPE ITP zone reached the interface shortly before 1.30 min (Fig. 9A and 9B). The stereoisomers leave the section containing the chiral selector within a short time (about 0.05 min, see broken line graphs in Fig. 9A and data of Fig. 9B and 9C). Both become concentrated to 6.84 mM and jointly form a migrating ITP zone of uniform conductivity (Fig. 9A). Initially, this ITP zone exhibits a non-uniform distribution of the stereoisomers. (-)-NPE is still migrating ahead of (+)-NPE (see 1.35 min distribution in Fig. 9A). With time and upon continued application of power, the interface between the two becomes broader due to diffusion (see 1.50 to 2.50 min time points of Fig. 9A). A uniform ITP zone with a concentration of 3.42 mM of each stereoisomer is eventually formed (data not shown). For both NPE

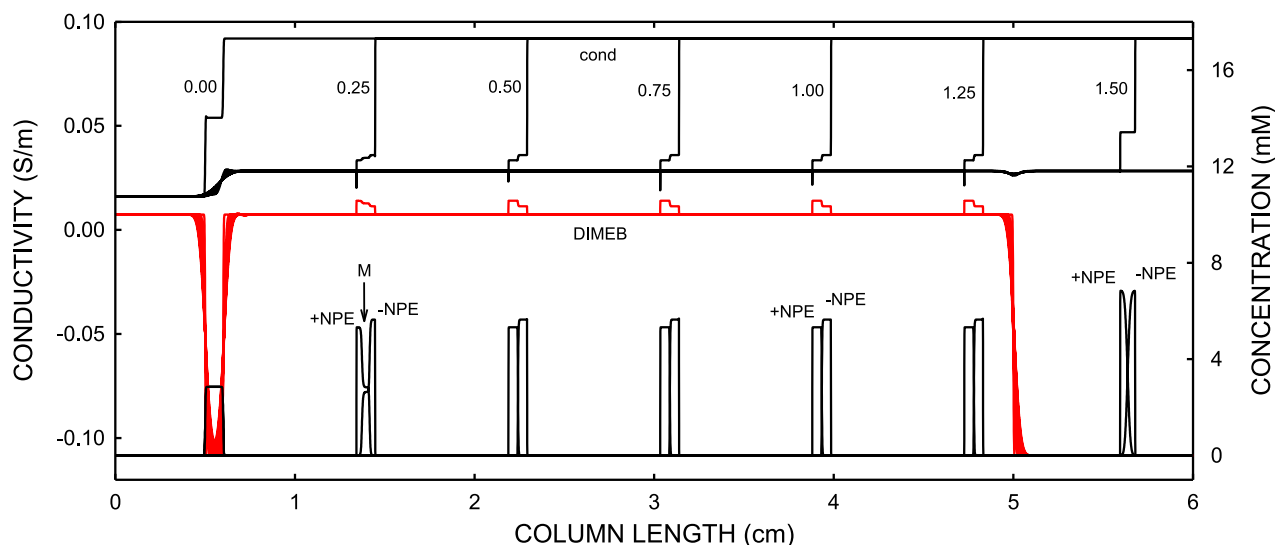


Figure 8. Computer predicted distributions of the NPE stereoisomers, DIMEB (red line graphs) and conductivity while the NPE ITP zones migrate within the leader zone containing DIMEB (up to 5 cm of column length, 0 to 1.25 min at 0.25 min interval) and thereafter (1.50 min). The two NPE stereoisomers were applied as hydrochlorides. Other conditions and key are identical as those of Figs. 1 and 2.

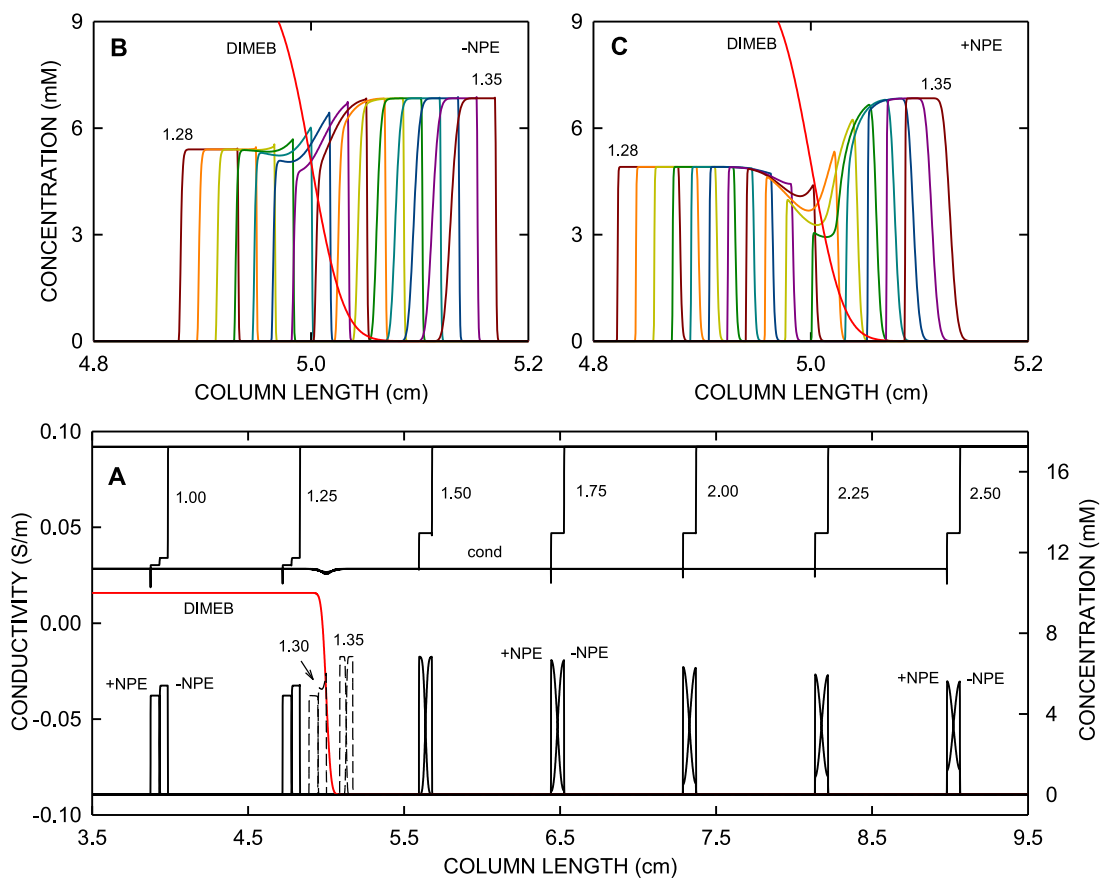


Figure 9. Computer predicted stereoisomer and conductivity distributions while and after the NPE ITP zones migrate across the end of immobilized DIMEB at 5 cm of column length. (A) NPE concentration profiles (lower graphs), DIMEB distribution (red line graph) and conductivity profiles (upper graphs) between 1.0 and 2.5 min (0.25 min interval) of power application. (B) (-)-NPE and (C) (+)-NPE concentration profiles in the transition region with vanishing DIMEB predicted for 1.280 to 1.350 min at an interval of 0.005 min. Other conditions and key as for Fig. 7.

analytes, the transition between the column segment with immobilized DIMEB to that without DIMEB is characterized with first a decrease in concentration followed by an increase to 6.84 mM (Fig. 9B and 9C, NPE profiles between 1.280 and 1.350 min at an interval of 0.005 min). The former effect is more pronounced

for (+)-NPE, the stereoisomer with stronger complexation. Comparable data were obtained for the case of free solution (Fig. 8, detailed profiles not shown). All these data suggest that analyte detection should occur shortly after the segment with the chiral selector.

4. Conclusions

For two NPE stereoisomers as model compounds, simulated zone patterns for the case of cationic ITP in free solution with DIMEB as selector were found to compare qualitatively well with those monitored experimentally with conductivity and absorbance detectors placed along coated capillaries. Stereoisomer separation was followed with an array of 8 conductivity detectors and occurs due to the presence of the selector. The technique used is ITP but the separation mechanism involved is the same as in electrokinetic chromatography. The separation of the NPE stereoisomers could also be simulated in presence of immobilized DIMEB, an approach that is reported here for the first time and remains to be experimentally validated. Compared to ITP in free solution, immobilization resulted in zone properties, including NPE concentrations, conductivity and pH, that were somewhat lower. The lower plateau concentrations of the analytes suggest that it is preferable to use a free rather than an immobilized neutral selector for ITP. Column preparation and the possibility for a rapid change of the separating medium are other advantages associated with a free selector. Finally, simulation provided insight into the migration behavior of ITP zones that migrate across a vanishing selector edge (end of selector zone) into the leader without chiral selector. In this process, NPE zones become concentrated and form a migrating ITP zone of uniform conductivity in which the two stereoisomers first migrate one after the other and then slowly become mixed due to diffusion. This is the first time that this process was elucidated and visualized. The simulation data reveal that highest sensitivity for analyte detection is obtained when the detector is placed immediately after the end of the selector zone. This is important for analyte detection with MS when the selector should not enter the detection system. Partial filling of the column with the selector is otherwise not an appealing approach for ITP. Simulation represents an elegant and efficient way to provide insight into the formation of transient analyte profiles, boundaries and zones in ITP with complexation, decomplexation and diffusional remixing outside the selector zone.

Author statement

Jitka Caslavská: Experimental work, data analysis, figure preparation, manuscript reviewing

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Wolfgang Thormann: Conceptualization, funding acquisition, simulations, manuscript writing

Declaration of Competing Interest

The authors have declared no conflict of interests.

Acknowledgments

Discussions with Dr. Peter C. Hauser and the technical assistance of Israel Joel Koenka provided during the generation of the conductivity array detector data are gratefully acknowledged. This work was partly supported by the Swiss National Science Foundation.

References

- [1] F.M. Everaerts, J.L. Beckers, Th.P.E.M. Verheggen, *Isotachophoresis*, Elsevier, Amsterdam, 1976.
- [2] W. Thormann, Principles of isotachophoresis and dynamics of the isotachophoretic separation of 2 components, *Sep. Sci. Technol.* 19 (1984) 455–467.
- [3] P. Boček, M. Deml, P. Gebauer, V. Dolník, *Analytical Isotachophoresis*, VCH, Weinheim, 1988.
- [4] P. Gebauer, P. Boček, Zone order in isotachophoresis - The concept of the zone existence diagram and its use in cationic systems, *J. Chromatogr.* 267 (1983) 49–65.
- [5] W. Thormann, R.A. Mosher, Theory of electrophoretic transport and separations: The study of electrophoretic boundaries and fundamental separation principles by computer simulation, *Adv. Electrophoresis* 2 (1988) 45–108.
- [6] R.A. Mosher, D.A. Saville, W. Thormann, *The Dynamics of Electrophoresis*, VCH Publishers, Weinheim, 1992.
- [7] T. Hirokawa, K. Watanabe, Y. Yokota, Y. Kiso, Bidirectional isotachophoresis 1. Verification of bidirectional isotachophoresis and simultaneous determination of anionic and cationic components, *J. Chromatogr.* 633 (1993) 251–259.
- [8] J. Caslavská, W. Thormann, Bidirectional isotachophoresis in open-tubular, untreated fused-silica capillaries, *J. Chromatogr. A* 772 (1997) 3–17.
- [9] G. Brouwer, G.A. Postema, Theory of separation in displacement electrophoresis, *J. Electrochem. Soc.* 117 (1970) 874–878.
- [10] P. Boček, M. Deml, B. Kaplanová, J. Janák, Analytical isotachophoresis concept of separation capacity, *J. Chromatogr.* 160 (1978) 1–9.
- [11] F.E.P. Mikkers, F.M. Everaerts, J.A.F. Peek, Isotachophoresis - Concepts of resolution, load-capacity and separation efficiency, *J. Chromatogr.* 168 (1979) 293–315.
- [12] F.E.P. Mikkers, F.M. Everaerts, J.A.F. Peek, Isotachophoresis - Concepts of resolution, load-capacity and separation efficiency 2. Experimental evaluation, *J. Chromatogr.* 168 (1979) 317–332.
- [13] W. Thormann, D. Arn, E. Schumacher, Monitoring of the isotachophoretic separation of 2 components with an array detector, *Sep. Sci. Technol.* 19 (1984) 995–1011.
- [14] T. Hirokawa, K. Nakahara, Y. Kiso, The separation process in isotachophoresis 2. Binary-mixtures and transient state models, *J. Chromatogr.* 463 (1989) 51–71.
- [15] T. Hirokawa, K. Nakahara, Y. Kiso, Separation process in isotachophoresis 3. Transient state models for a 3-component system, *J. Chromatogr.* 470 (1989) 21–41.
- [16] P. Boček, I. Miedziak, M. Deml, J. Janák, Use of complex-formation equilibria in analytical isotachophoresis of strong electrolyte ions - separation of halides and sulfates, *J. Chromatogr.* 137 (1977) 83–91.
- [17] I. Nukatsuka, M. Taga, H. Yoshida, Separation of lanthanides by capillary-tube isotachophoresis using complex-forming equilibria, *J. Chromatogr.* 205 (1981) 95–102.
- [18] I. Jelínek, J. Snopek, E. Smolková-Keulemansová, Use of cyclodextrins in isotachophoresis 2. Alpha-cyclodextrins, beta-cyclodextrins and gamma-cyclodextrins as leading electrolyte additives for the separation of ortho-substituted, meta-substituted and para-substituted halogenobenzoic acids, *J. Chromatogr.* 411 (1987) 153–159.
- [19] P. Gebauer, P. Boček, M. Deml, J. Janák, Isotachophoresis of kinetically labile complexes, *J. Chromatogr.* 199 (1980) 81–94.
- [20] J. Snopek, I. Jelínek, E. Smolková-Keulemansová, Chiral separation by analytical electromigration methods, *J. Chromatogr.* 609 (1992) 1–17.
- [21] S. Fanali, Identification of chiral drug isomers by capillary electrophoresis, *J. Chromatogr. A* 735 (1996) 77–121.
- [22] B. Chankvetadze, *Capillary Electrophoresis in Chiral Analysis*, John Wiley & Sons, Chichester, 1997.
- [23] B. Chankvetadze, Contemporary theory of enantioseparations in capillary electrophoresis, *J. Chromatogr. A* 1567 (2018) 2–25.
- [24] S. Zaugg, W. Thormann, Enantioselective determination of drugs in body fluids by capillary electrophoresis, *J. Chromatogr. A* 875 (2000) 27–41.
- [25] J. Caslavská, W. Thormann, Stereoselective determination of drugs and metabolites in body fluids, tissues and microsomal preparations by capillary electrophoresis (2000–2010), *J. Chromatogr. A* 1218 (2011) 588–601.
- [26] J. Snopek, I. Jelínek, E. Smolková-Keulemansová, Use of cyclodextrins in isotachophoresis 4. The influence of cyclodextrins on the chiral resolution of ephedrine alkaloid enantiomers, *J. Chromatogr.* 438 (1988) 211–218.
- [27] I. Jelínek, J. Dohnal, J. Snopek, E. Smolková-Keulemansová, Use of cyclodextrins in isotachophoresis 7. Resolution of structurally related and chiral phenothiazines, *J. Chromatogr.* 464 (1989) 139–147.
- [28] M. Lanz, J. Caslavská, W. Thormann, Enantiomeric separation of methadone by cyclodextrin-based capillary and recycling isotachophoresis, *Electrophoresis* 19 (1998) 1081–1090.
- [29] P. Hoffmann, H. Wagner, G. Weber, M. Lanz, J. Caslavská, W. Thormann, Separation and purification of methadone enantiomers by continuous- and interval-flow electrophoresis, *Anal. Chem.* 71 (1999) 1840–1850.
- [30] P. Kubačák, P. Mikuš, I. Valášková, E. Havránek, Chiral separation of alkylamine antihistamines in pharmaceuticals by capillary isotachophoresis with charged cyclodextrin, *Drug Dev. Ind. Pharm.* 33 (2007) 1199–1204.
- [31] S. Zaugg, J. Caslavská, R. Theurillat, W. Thormann, Characterization of the stereoselective metabolism of thiopental and its metabolite pentobarbital via analysis of their enantiomers in human plasma by capillary electrophoresis, *J. Chromatogr. A* 838 (1999) 237–249.
- [32] D. Kanišky, E. Šimuničová, E. Olvecká, A. Ferancová, Separations of enantiomers by preparative capillary isotachophoresis, *Electrophoresis* 20 (1999) 2786–2793.
- [33] E. Olvecká, M. Masar, D. Kanišky, M. Jöhnc, B. Stanislawski, Isotachophoresis separations of enantiomers on a planar chip with coupled separation channels, *Electrophoresis* 22 (2001) 3347–3353.
- [34] D.A. Jayawickrama, J.V. Sweedler, Chiral separation of nanomole amounts of alprenolol with cITP/NMR, *Anal. Bioanal. Chem.* 378 (2004) 1528–1535.

- [35] C.J. Jones, C.K. Larive, Microcoil NMR Study of the interactions between doxepin, beta-cyclodextrin and acetate during capillary isotachopheresis, *Anal. Chem.* 84 (2012) 7099–7106.
- [36] S. Mikkonen, J. Caslavská, P. Gebauer, W. Thormann, Inverse cationic ITP for separation of methadone enantiomers with sulfated beta-cyclodextrin as chiral selector, *Electrophoresis* 40 (2019) 659–667.
- [37] W. Thormann, J. Caslavská, M.C. Breadmore, R.A. Mosher, Dynamic computer simulations of electrophoresis: Three decades of active research, *Electrophoresis* 30 (2009) S16–S26.
- [38] W. Thormann, M.C. Breadmore, J. Caslavská, R.A. Mosher, Dynamic computer simulations of electrophoresis: a versatile research and teaching tool, *Electrophoresis* 31 (2010) 726–754.
- [39] W. Thormann, C.-X. Zhang, J. Caslavská, P. Gebauer, R.A. Mosher, Modeling of the impact of ionic strength on the electroosmotic flow in capillary electrophoresis with uniform and discontinuous buffer systems, *Anal. Chem.* 70 (1998) 549–562.
- [40] V. Hruška, M. Jaroš, B. Gaš, Simul 5 – Free dynamic simulator of electrophoresis, *Electrophoresis* 27 (2006) 984–991.
- [41] M. Bercovici, S.K. Lele, J.G. Santiago, Open source simulation tool for electrophoretic stacking, focusing, and separation, *J. Chromatogr. A* 1216 (2009) 1008–1018.
- [42] R.A. Mosher, M.C. Breadmore, W. Thormann, High-resolution electrophoretic simulations: performance characteristics of one-dimensional simulators, *Electrophoresis* 32 (2011) 532–541.
- [43] S. Mikkonen, H. Elkstrom, W. Thormann, High-resolution dynamic computer simulation of electrophoresis using a multiphysics software platform, *J. Chromatogr. A* 1532 (2018) 216–222.
- [44] E. Dubrovčáková, B. Gaš, J. Vacík, E. Smolková-Keulemansová, Electromigration in systems with additives in background electrolytes 1. Addition of the neutral complexing agent, *J. Chromatogr.* 623 (1992) 337–344.
- [45] M.C. Breadmore, H.Y. Kwan, J. Caslavská, W. Thormann, Dynamic high-resolution computer simulation of electrophoretic enantiomer separations with neutral cyclodextrins as chiral selectors, *Electrophoresis* 33 (2012) 958–969.
- [46] V. Hruška, M. Beneš, J. Svobodová, I. Zusková, B. Gaš, Simulation of the effects of complex-formation equilibria in electrophoresis: I. Mathematical model, *Electrophoresis* 33 (2012) 938–947.
- [47] J. Svobodová, M. Beneš, V. Hruška, K. Ušelová, B. Gaš, Simulation of the effects of complex-formation equilibria in electrophoresis: II. Experimental verification, *Electrophoresis* 33 (2012) 948–957.
- [48] J. Caslavská, M.C. Breadmore, W. Thormann, Dynamic high-resolution computer simulation of isotachophoretic enantiomer separation and zone stability, *Electrophoresis* 35 (2014) 625–637.
- [49] J. Caslavská, W. Thormann, Contemporary chiral simulators for capillary zone electrophoresis, *Electrophoresis* 41 (2020) 502–513.
- [50] J. Caslavská, I.J. Koenka, P.C. Hauser, W. Thormann, Validation of CE modelling with a contactless conductivity array detector, *Electrophoresis* 37 (2016) 699–710.
- [51] P. Boček, P. Gebauer, M. Deml, Migration behavior of the hydrogen-ion and its role in isotachopheresis of cations, *J. Chromatogr.* 217 (1981) 209–224.
- [52] V.K. Almeida, C.K. Larive, Insights into cyclodextrin interactions during sample stacking using capillary isotachopheresis with on-line microcoil NMR detection, *Magn. Reson. Chem.* 43 (2005) 755–761.
- [53] M. Dovhunová, M. Malý, P. Dubský, G.S. Gerlero, P.A. Kler, Generalized model of the linear theory of electromigration and its application to electrokinetic chromatography: Capillary zone electrophoretic systems with complex-forming equilibria, *J. Chromatogr. A* 1610 (2020) 460595.
- [54] J. Caslavská, R.A. Mosher, W. Thormann, Impact of Taylor-Aris diffusivity on analyte and system zone dispersion in CZE assessed by computer simulation and experimental validation, *Electrophoresis* 36 (2015) 1529–1538.
- [55] W. Thormann, J. Caslavská, R.A. Mosher, Computer simulation of electrophoretic aspects of enantiomer migration and separation in capillary electrochromatography with a neutral selector, *Electrophoresis* 36 (2015) 773–783.
- [56] V. Schurig, D. Wistuba, Recent innovations in enantiomeric separation by electrochromatography utilizing modified cyclodextrins as stationary phases, *Electrophoresis* 20 (1999) 2313–2328.
- [57] D. Wistuba, V. Schurig, Cyclodextrin-mediated enantioseparations by capillary electrochromatography, *Methods Mol. Biol.* 970 (2013) 505–523.
- [58] M. Riesová, J. Svobodová, Z. Tošner, M. Beneš, E. Tesařová, B. Gaš, Complexation of buffer constituents with neutral complexation agents: Part I. Impact on common buffer properties, *Anal. Chem.* 85 (2013) 8518–8525.